## AMENDMENT TO THE CLAIMS:

Please amend the claims as follows

Please cancel claims 7, 9, 32, 44, 45, 50 to 61, 63 to 74, 100 to 103, 109 and 130, without prejudice or disclaimer.

This listing of claims will replace all prior versions and listings of claims in the application: Listing of Claims:

Claim 1 (currently amended): A method for producing a recombinant antibody or antigen binding fragment thereof with improved yield from a host cell, comprising:

- (i) providing a nucleic acid encoding a modified-non-human antibody or antigen binding fragment thereof made by a method comprising:
  - (a) aligning a hypervariable region (HVR1) and/or a hypervariable region 2 (HVR2) of a heavy chain variable domain of a non-human antibody or antigen binding fragment thereof to corresponding HVR1 and/or HVR2 sequences of human subgroup variable domain consensus sequences;
  - (b) selecting a human subgroup variable domain consensus sequence that has a HVR1 and/or HVR2 amino acid sequence with the most sequence identity with the non-human HVR1 sequence and/or the non-human HVR2 sequence:
  - (c) identifying at least one amino acid position in at least one framework region (FR) of the selected human subgroup variable domain consensus sequence that has a different amino acid residue than that of a corresponding position in a FR of the variable domain of the non-human antibody or antigen binding fragment thereof; and
  - (d) <u>substituting at least modifying</u> one amino acid <u>in the FR</u> at the corresponding position of the non-human variable domain of the antibody or <u>the</u> antigen binding fragment <u>thereof identified in step (i)(c)</u> with the corresponding to be the same as the different human-amino acid <u>from the selected subgroup consensus sequence in step (b)</u> residue identified in (e) to form a <u>substituted modified FR region</u> in the non-human variable domain of the antibody or antigen binding fragment <u>thereof</u>; and
- (ii) expressing the <u>substituted</u> <del>modified</del>-non-human antibody or antigen binding fragment thereof in the host cell.

wherein the <u>substituted</u> modified non-human antibody or antigen binding fragment <u>thereof</u> having the at least one substitution in the FR has improved yield in a cell or a cell culture as compared to the corresponding <u>unsubstituted</u> <del>unmodified</del> antibody or antigen binding fragment thereof.

Claim 2 (currently amended): The method according to claim 1, wherein the non-human antibody or antigen binding fragment thereof to be substituted modified is selected from the group consisting of a humanized antibody, a chimeric antibody, a monoclonal antibody, a multispecific antibody, a diabody, or an antibody generated by phage display.

Claim 3 (currently amended): The method according to claim 2, wherein the non-human antigen binding fragment thereof is a Fab fragment, F(ab')<sub>2</sub> fragment, scFV fragment, or sc(Fv)<sub>2</sub> fragment, a single arm antibody or single chain antibody.

Claim 4 (currently amended): The method according to claim 1, wherein the non-human antibody is an anti-vascular endothelial growth factor (VEGF) anti-VEGF antibody.

Claim 5 (previously presented): The method according to claim 4, wherein the non-human antibody is a humanized antibody.

Claims 6 and 7 (canceled)

Claim 8 (currently amended): The method of claim 1, wherein the host-cell-comprises an expression vector comprising the nucleic acid encoding the <u>substituted recombinant modified</u>-nonhuman antibody or antigen binding fragment thereof is contained in an expression vector.

Claim 9 (canceled)

Claim 10 (previously presented): The method according to claim 1, wherein the host cell is a prokaryotic host cell.

Claim 11 (previously presented): The method according to claim 1, wherein the host cell is a mammalian cell.

Claim 12 (currently amended): The method according to claim 1, further comprising isolating the expressed non-human heavy chain variable domain having a-modified substituted FR region-or the-modified substituted non-human light chain variable domain having a-modified substituted FR-region from the cell or cell culture.

Claim 13 (currently amended): The method according to claim 12, wherein the non-human variable domain is a heavy chain variable domain and the HVR1 amino acid sequence of the heavy chain variable domain of <a href="mailto:said">said</a> [[the]] antibody or antigen binding fragment <a href="mailto:thereof">thereof</a> is GYTFTNYGIN (SEQ ID NO: 14) or GYDFTHYGMN (SEQ ID NO:18).

Claim 14 (currently amended): The method according to claim 1, wherein the non-human framework region to be modified substituted is selected from the group consisting of a FR1, a FR2, a FR3, a FR4 and a mixture thereof.

Claim 15 (currently amended): The method according to claim 14, wherein the human subgroup variable domain consensus sequence comprises a variable domain FR1 sequence that is with a sequence-selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3.

Claim 16 (currently amended): The method according to claim 1, wherein the yield of the non-human antibody or antigen binding fragment <u>thereof</u> comprising the <u>modified substituted</u> FR is improved at least 2 fold compared to the corresponding <del>unmodified <u>unsubstituted</u></del> antibody or antigen binding fragment <u>thereof</u>.

Claim 17 (currently amended): The method according to claim 16, wherein the yield of the non-human antibody or antigen binding fragment thereof comprising the modified substituted FR is

improved at least 2 fold to 16 fold compared to the corresponding-unmodified unsubstituted antibody or antigen binding fragment thereof,

Claim 18 (currently amended): The method of claim 1, wherein in step (d) two, three, four, five, six or seven amino acid positions in the non-human FR are <u>substituted-modified</u>.

Claim 19 (currently amended): The method of claim 1, wherein the non-human antibody or antigen binding fragment thereof is an anti-vascular endothelial growth factor (VEGF) a VEGF antibody or antigen binding fragment thereof comprising a heavy chain variable domain FR1 sequence of SEQ ID NO:3, wherein and the FR is a heavy chain variable domain FR1-and-one of the amino acid positions is position 6 or position 23 or both, and the other position is selected from the group consisting of position 1, 11, 13, 18, 19, and a mixture thereof, wherein the corresponding amino acid residues are found in the heavy chain FR1 consensus sequence of SEQ ID NO:1.

Claim 20 (currently amended): The method of claim 19, wherein amino acid positions 6 and 23 are substituted modified.

Claim 21 (currently amended): The method of claim 19, wherein the amino acid positions at positions 1, 6, 11, 13, 18, 19, and 23 of the heavy chain FR1 are substituted modified.

Claim 22 (currently amended): The method of claim 1, wherein at least two one-but not all of the different amino acid positions in the non-human FR are substituted modified.

Claim 23 (currently amended): The method of claim 22, wherein the modified substituted FR is FR1, FR2, or FR3.

Claim 24 (currently amended): The method of claim 1, wherein-at-least-one-but-not all of the amino acid positions that have a different amino acid as compared to the human consensus sequence in all framework regions (FRs) of the non-human variable region are substituted-modified.

Claim 25 (currently amended): A method for preparing a humanized antibody or an antigen binding fragment <u>thereof</u> having an improved folding efficiency and yield when expressed in a host cell, comprising:

- (a) preparing a humanized antibody or antigen binding fragment thereof comprising a variable domain comprising at least one-modified <u>substituted</u> framework <u>region</u> (FR) sequence, wherein the variable domain is made by a method comprising:
  - (i) aligning a hypervariable region (HVR1) and/or a hypervariable region 2 (HVR2) of a variable domain of a non-human antibody or antigen binding fragment thereof to corresponding HVR1 and/or HVR2 sequences of human subgroup variable domain consensus sequences;
  - (ii) selecting a human subgroup variable domain consensus sequence that has a HVR1 and/or HVR2 amino acid sequence with the most sequence identity with the non-human HVR1 sequence and/or the non-human HVR2 sequence;
  - (iii) identifying at least one amino acid position in at least one framework region (FR) of the human subgroup variable domain consensus sequence selected in step (ii) that has a different amino acid residue than that of a corresponding position in a FR of the variable domain or antigen binding fragment thereof of the non-human antibody; and
  - (iv) <u>substituting modifying</u> one amino acid <u>in the FR</u> at the eorresponding <u>position</u> of the non-human variable domain or antigen binding fragment <u>thereof</u> <u>identified in step (a)(ii)</u> with the corresponding of the antibody to be the same as the <u>different human</u> amino acid <u>from the selected subgroup consensus sequence in step (a)(iii)</u> residue identified in (e) to form a modified <u>substituted</u> FR region in the non-human variable domain or antigen binding fragment <u>thereof</u> of the antibody,
  - wherein the <u>substitution modification</u> results in an antibody or antigen binding fragment <u>thereof</u> having an improved folding efficiency and yield when expressed in the host cell, and
- (b) expressing the-modified <u>substituted</u> humanized antibody or-modified <u>substituted</u> antigen binding fragment <u>thereof</u> in the host cell.

Claims 26 to 27 (canceled)

Claim 28 (currently amended): The method according to claim 25 wherein the non-human variable domain is a heavy chain variable domain and the HVR1 amino acid sequence of a heavy chain variable domain of the non-human antibody or antigen binding fragment <u>thereof</u> is GYTFTNYGIN (SEQ ID NO: 14) or GYDFTHYGMN (SEQ ID NO:18).

Claim 29 (previously presented): The method according to claim 25, wherein the FR is selected from the group consisting of a FR1, a FR2, a FR3, a FR4 and a mixture thereof.

Claim 30 (currently amended): The method according to claim 29, wherein the human subgroup variable domain consensus sequence comprises a heavy chain variable domain FR1 sequence that is with a sequence-selected from the group consisting of SEQ. ID NO: 1, SEQ ID NO: 2 and SEQ ID NO: 3.

Claim 31 (currently amended): The method of claim 25, wherein at least one but not all of the different amino acid positions in the non-human FR identified in (a)(iii) are modified substituted.

Claim 32 (canceled)

Claim 33 (currently amended): The method of claim 25, wherein the host cell comprises an expression vector comprising the nucleic acid encoding the <u>substituted recombinant modified-non-</u> human antibody or antigen binding fragment thereof is contained in an expression vector.

Claim 34 (currently amended): The method of claim 25 [[33]], wherein the nucleic acid further comprises a sequence encoding a constant domain connected to the nucleic acid encoding the modified substituted non-human antibody or antigen binding fragment thereof to form a nucleic acid encoding a full-length heavy or light chain.

Claim 35 (canceled)

Claim 36 (previously presented): The method according claim 25, wherein the host cell is a prokaryotic host cell.

Claim 37 (previously presented): The method according to claim 25, wherein the host cell is a mammalian cell.

Claim 38 (currently amended): A method for improving the yield of an assembled nonhuman monoclonal antibody or antigen binding fragment thereof in a host cell, comprising:

- (a) aligning a hypervariable region 1 (HVR1) and/or a hypervariable 2 (HVR2) sequence of a heavy chain variable domain of the non-human monoclonal antibody to corresponding HVR1 and/or HVR2 sequences of human subgroup heavy chain variable domain consensus sequences,
- (b) selecting a human subgroup heavy chain variable domain consensus sequence that has the HVR1 and/or HVR2 amino acid sequence with the most sequence identity with the HVR1 and/or the HVR2 sequence of the heavy chain variable domain of the non human monoclonal antibody,
- (c) <u>substituting modifying-at least one but not all-amino acid position positions-in at least</u> one framework <u>region</u> (FR) of the non-human monoclonal antibody heavy chain variable domain <u>for [[tol]]</u> an amino acid residue found at a corresponding position <u>from [[of]]</u> the selected human subgroup heavy chain variable domain consensus sequence to form at least one-<u>modified substituted</u> FR, wherein the non-human monoclonal antibody or antigen binding fragment <u>thereof</u> with the <u>modified substituted</u> FR has improved folding efficiency and yield, in cell culture compared to the folding efficiency and yield of a corresponding-<del>unmodified unsubstituted</del> antibody or antigen binding fragment <u>thereof</u>; and
- (d) expressing the non-human monoclonal antibody or antigen binding fragment thereof comprising the modified substituted FR in the host cell.

Claim 39 (currently amended): A method for improving the yield of a recombinant antibody or antigen binding fragment thereof expressed in a host cell, comprising:

(a) selecting a human subgroup variable domain consensus sequence by aligning a hypervariable region 1 (HVR1) and/or a hypervariable 2 (HVR2) sequence of a variable domain of

a non-human antibody or antigen binding fragment thereof to corresponding HVR1 and/or HVR2 sequences of human subgroup variable domain consensus sequences, and selecting the human subgroup variable domain consensus sequence that has the HVR1 and/or HVR2 amino acid sequence with the most sequence identity with the HVR1 and/or HVR2 sequence of the variable domain of the non-human antibody or antigen binding fragment thereof. [fand]]

(b) <u>substituting</u> modifying at least one but not all amino acid <u>residue</u> residues in the framework <u>region</u> (FR) of the variable domain of the non-human antibody or antigen binding fragment <u>thereof for the corresponding amino acid from such that the modified FR has at least 50% sequence identity to the corresponding FR amino acid sequence of the selected human subgroup variable domain consensus sequence to form a modified FR.</u>

wherein the amino acid residues in the framework (FR) are modified to the amino acid residue of the corresponding human subgroup variable domain consensus sequence amino acid,

wherein [[and]] the recombinant antibody or antigen binding fragment thereof with the modified substituted FR has improved folding efficiency and yield [[,]] in cell culture compared to the folding efficiency and yield of a corresponding-unmodified unsubstituted antibody or antigen binding fragment thereof; and

(c) expressing the <u>recombinant</u> antibody or antigen binding fragment <u>thereof</u> with the <u>modified</u> <u>substituted</u> FR in the host cell and recovering the antibody or antigen binding fragment thereof with the <u>modified</u> substituted FR from the host cell.

Claim 40 (currently amended): The method according to claim 39, wherein the variable domain is a heavy chain variable domain and the HVR1 amino acid sequence of a heavy chain variable domain of the <u>non-human</u> antibody or antigen binding fragment <u>thereof</u> is GYTFTNYGIN (SEQ ID NO: 14) or GYDFTHYGMN (SEQ ID NO: 18).

Claim 41 (currently amended): The method of claim 39, wherein at least two but not all amino acid positions that have a different amino acid in at least one FR are substituted with amino acids in the corresponding position of the selected human subgroup consensus sequence.

Claim 42 (currently amended): The method of claim 41, wherein the antibody or antibody binding fragment thereof is an anti-vascular endothelial growth factor (VEGF) a-VEGF-antibody or antibody binding fragment thereof comprising a heavy chain variable domain FR1 comprising the amino acid sequence of SEQ ID NO:3 and amino acid positions 6 and 23 of SEQ ID NO:3 heavy chain FR1-are-modified substituted with the corresponding amino acid found in SEQ ID NO:1.

Claim 43 (currently amended): The method of claim 42, wherein amino acid positions 1, 6, 11, 13, 18, 19 and 23 of the heavy chain FR1 of SEQ ID NO:3 are-modified substituted.

Claims 44 to 45 (canceled)

Claim 46 (previously presented): The method according to claim 38, wherein the host cell is a prokaryotic host cell.

Claim 47 (previously presented): The method according to claim 38, wherein the host cell is a mammalian cell.

Claim 48 (currently amended): The method according to claim 39, wherein the step [[of]] (b) comprises substituting at least modifying one but not all amino acid residue residues in all of the FRs of the variable domain of the non-human antibody or antigen binding fragment thereof with the corresponding amino acid residues [[of]] from the selected corresponding-human variable domain subgroup-variable domain consensus sequence.

Claim 49 (previously presented): The method according to claim 38, wherein the framework region sequence is selected from the group consisting of FR1, FR2, FR3, FR4 and a mixture thereof.

Claims 50 to 95 (canceled)

Claim 96 (currently amended): A method for improving the yield of antibody or antigen binding fragment thereof in a host cell or cell culture, comprising:

a) expressing a nucleic acid encoding a variable domain of a non-human antibody or antigen binding fragment thereof comprising at least one modified substituted framework region (FR) in the host cell, wherein the modified substituted FR has: [[(i)]] a substitution of at least one but not all amino acids in the at least one FR with a different amino acid, or (ii) a deletion of at least one but not all amino acids in the FR.

wherein the amino acid residue or residues to be substituted or deleted is determined by aligning a hypervariable region 1 (HVR1) and/or hypervariable region 2 (HVR2) sequence of the non-human variable domain of the antibody or antigen binding fragment thereof to corresponding HVR1 and/or HVR2 sequences of human subgroup variable domain consensus sequences, and selecting the amino acid found at the corresponding FR position of the human subgroup variable domain consensus sequence that has a HVR1 and/or HVR2 amino acid sequence with the most sequence identity with the HVR1 and/or HVR2 sequence of the non-human variable domain of the recombinant antibody or antigen binding fragment thereof, and

b) recovering the <u>recombinant</u> antibody or antigen binding fragment <u>thereof</u> comprising the non-human variable domain comprising the <u>modified substituted</u> FR from the host cell, wherein the <u>modified substituted</u> antibody or antigen binding fragment <u>thereof</u> has improved folding efficiency and yield in the cell or cell culture as compared to the folding efficiency and yield of an <u>unmodified</u> <u>unsubstituted</u> antibody or antigen binding fragment <u>thereof</u>.

Claim 97 (previously presented): The method according to claim 96, wherein:

(a) the nucleic acid is contained in an expression vector, (b) the nucleic acid is operably linked to a promoter, (c) the method of (a) or (b), wherein the nucleic acid further comprises a heat stable enterotoxin sequence that can direct secretion to the periplasm, or (d) the method of any of (a) to (c), wherein the nucleic acid further comprises a terminator sequence.

Claim 98 (previously presented): The method according to claim 96, wherein the host cell is a prokaryotic host cell.

Claim 99 (previously presented): The method according to claim 96, wherein the host cell is a eukaryotic host cell.

Claims 100 to 103 (canceled)

Claim 104 (currently amended): A method for improving the yield of a non-human antibody, or antigen binding fragments thereof, in a host cell or cell culture comprising:

- (a) comparing a hypervariable region 1 (HVR1) and/or hypervariable region 2 (HVR2) amino acid sequence of a heavy chain variable domain of the non-human antibody or antigen binding fragment thereof to a corresponding HVR1 and/or HVR2 amino acid sequence of each human subgroup heavy chain variable domain consensus sequence and selecting the human subgroup heavy chain variable domain consensus sequence that has the most sequence identity with the HVR1 and/or HVR2 sequence of the heavy chain variable domain of the non-human antibody or antigen binding fragment thereof;
- (b) identifying at least one amino acid position in at least one framework region (FR) in the heavy chain variable domain of the non-human antibody or antigen binding fragment thereof, wherein the FR is selected from the group consisting of a FR1, a FR2, a FR3, a FR4 and a mixture thereof, wherein the amino acid position has a different amino acid than the amino acid at a corresponding position of the selected human subgroup heavy chain variable domain consensus sequence; and
- (c) <u>substituting said modifying or deleting at least one but not all of the amino acid positions</u> identified in step (b), <u>with wherein the modification or deletion is with the amino acid in the</u> corresponding <u>amino acid from position of</u> the selected human heavy chain subgroup variable domain consensus sequence, to form a variable domain with a <u>modified</u> <u>substituted</u> FR; and
- (d) expressing the <u>non-human</u> antibody or antigen binding fragment <u>thereof</u> comprising the heavy chain variable domain with the <u>modified substituted</u> FR in the host cell or cell culture, and
- (e) recovering the antibody or antigen binding fragment thereof from the host cell or cell culture.

wherein the <u>non-human</u> antibody or antigen binding fragment <u>thereof</u> with the <u>modified</u> substituted FR has improved yield in the host cell or cell culture compared to the folding efficiency

and yield of a corresponding-unmodified unsubstituted non-human antibody or antigen binding fragment thereof.

Claim 105 (previously presented): The method according to claim 104, wherein the nonhuman antibody is selected from the group consisting of a humanized antibody, a chimeric antibody, a monoclonal antibody, a multispecific antibody, a diabody, or an antibody generated by phage display.

Claim 106 (currently amended): The method according to claim 105, wherein the nonhuman antigen binding fragment thereof is a Fab fragment, F(ab')<sub>2</sub> fragment, scFV fragment, or sc(Fv)<sub>2</sub> fragment, single arm antibody, or single chain antibody.

Claim 107 (previously presented): The method according to claim 104, wherein the nonhuman antibody is an anti-VEGF antibody.

Claim 108 (previously presented): The method according to claim 107, wherein the nonhuman antibody is a humanized antibody.

Claim 109 (canceled)

Claim 110 (currently amended): The method of claim [[109]] 104, wherein the variable domain domain encoding nucleic acid further comprises a nucleic acid encoding a constant region domain, and the constant region domain domain encoding nucleic acid is connected to the nucleic acid encoding the variable domain with the modified substituted FR to form a nucleic acid encoding a-variant full-length heavy or light chain.

Claim 111 (currently amended): The method of claim [[109]] 104, wherein the host cell comprises modified nucleic acid is comprised within an expression vector comprising the nucleic acid sequence that encodes the antibody or antigen binding fragment thereof with the substitute FR.

Claim 112 (currently amended): The method of claim 111, further comprising culturing [[a]]

the host cell comprising the expression vector-or the modified nucleic acid under conditions wherein
the recombinant antibody or antigen binding fragment thereof is chains are expressed; and
recovering said antibody or antigen binding fragment thereof a full length heavy or light chain or
both-from the cell or cell culture.

Claim 113 (original): The method according to claim 112, wherein the host cell is a prokaryotic host cell.

Claim 114 (original): The method according to claim 112, wherein the host cell is a mammalian cell.

Claim 115 (canceled)

Claim 116 (previously presented): The method according to claim 104, wherein the variable domain is a heavy chain variable domain and the HVR1 amino acid sequence is GYTFTNYGIN (SEQ ID NO: 14) or GYDFTHYGMN (SEQ ID NO:18).

Claim 117 (previously presented): The method according to claim 104, wherein the framework region is selected from the group consisting of FR1, FR2, FR3, and a mixture thereof.

Claim 118 (currently amended): The method according to claim 117, wherein the human subgroup FR consensus sequence is a heavy chain FR1 sequence that is with a sequence-selected from the group consisting of SEO ID NO: 1, SEO ID NO: 2 and SEO ID NO: 3.

Claim 119 (currently amended): The method according to claim 104, wherein the yield of the antibody or antigen binding fragment thereof with the modified substituted FR is improved at least 2 fold compared to the corresponding unmodified unsubstituted antibody or antigen binding fragment thereof.

Claim 120 (currently amended): The method according to claim 119, wherein the yield of the antibody or antigen binding fragment thereof with the modified substituted FR is improved at least 2 fold to 16 fold compared to the corresponding-unmodified unsubstituted antibody or antigen binding fragment thereof.

Claim 121 (currently amended): The method of claim 104, wherein at least two-but not all of the identified amino acid positions in at least one FR of the non-human antibody or antigen binding fragment thereof are: [[(i)]] substituted with amino acids in the corresponding position of the selected human subgroup consensus sequence, or (ii) deleted.

Claim 122 (currently amended): The method of claim 121, wherein the non-human antibody or antigen binding fragment thereof is an anti-vascular endothelial growth factor (VEGF) a VEGF antibody or antigen binding fragment thereof that comprises a heavy chain variable domain FR1 comprising the amino acid sequence of SEQ ID NO:3, and the FR is a heavy chain FR1 and one of the identified amino acid positions is position 6 or position 23 or both, and the other position is selected from the group consisting of position 1, 11, 13, 18, 19, and a mixture thereof of SEQ ID NO:3 are substituted with the corresponding amino acids from SEQ ID NO:1.

Claim 123 (currently amended): The method of claim 122 wherein amino acid positions 6 and 23 are substituted with the corresponding amino acids from SEQ ID NO:1.

Claim 124 (currently amended): The method of claim 122, wherein all of the amino acid positions at positions at position [[.]] 1, 6, 11, 13, 18, 19 [[,]] and 23 of the heavy chain FR1 are substituted with the corresponding amino acids from SEQ ID NO:1.

Claim 125 (currently amended): The method of claim 104, wherein at least three but not all of the identified amino acid positions in a FR are [[: (i)]] substituted with the amino acid in the corresponding position in the selected human subgroup consensus sequence—or (ii) deleted.

Claim 126 (previously presented): The method of claim 125, wherein the FR is a FR1, a FR2, or a FR3.

Claim 127 (currently amended): The method of claim 104, wherein at least four but not all of the identified amino acid positions in all FR are; (i) substituted with the amino acid in the corresponding position in the selected subgroup consensus sequence, or (ii) deleted.

Claim 128 (canceled)

Claim 129 (currently amended): The method of claim 104 further comprising:

- (a) comparing a hypervariable region 1 (HVR1) and/or hypervariable region 2 (HVR2) amino acid sequence of a light chain variable domain of a non-human antibody or antigen binding fragment thereof to a corresponding HVR1 and/or HVR2 amino acid sequence of a human subgroup light chain variable domain consensus sequence and selecting the human subgroup light chain variable domain consensus sequence that has the most sequence identity with the HVR1 and/or HVR2 sequence of the non-human light chain variable domain;
- (b) identifying at least one amino acid position in at least one FR in the non-human light chain variable domain wherein the FR is selected from the group consisting of a FR1, a FR2, a FR3, a FR4 and a mixture thereof, wherein the amino acid position has a different amino acid than the amino acid at a corresponding position of the selected human subgroup light chain variable domain consensus sequence; and
- (c) (i) <u>substituting</u> modifying the at least one but not all of the non-human amino acid in the <u>FR</u> of the non-human positions identified in step (b) with the amino acid in the corresponding position of the selected human subgroup light chain-variable domain of the antibody or the antigen binding fragment thereof identified in step (b) with the corresponding amino acid from the selected <u>subgroup</u> consensus sequence to form a modified light chain-variable domain with a modified FR, or (ii) deleting the at least one but not all of the non-human amino acid positions identified in step (b) <u>substituted FR</u> in the non-human variable domain of the antibody or antigen binding fragment thereof.

Claim 130 (canceled)

Claim 131 (previously presented): The method of claim 1, wherein the host cell is a prokaryotic cell or a eukaryotic cell.

Claim 132 (previously presented): The method of claim 131, wherein the host cell is a filamentous fungi or yeast cell, an insect cell, a mammalian cell or a bacterial cell.

Claim 133 (currently amended): The method of claim 132, wherein the host cell is an Archaebacteria or a Eubacteria, or a Gram-negative or a Gram-positive bacteria-organism.